CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 21-334 and 21-085/S-010

APPROVED DRAFT LABELING

AVELOX®

(moxifloxacin hydrochloride) Tablets

DESCRIPTION

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AVELOX® (moxifloxacin hydrochloride) is a synthetic broad spectrum antibacterial agent for oral administration. Moxifloxacin, a fluoroguinolone, is available as the monohydrochloride salt of 1-cyclopropyl-7-[(S,S)-2,8-diazabicyclo[4.3.0]non-8-yl]-6-fluoro-8-methoxy-1,4-dihydro-4-oxo-3-quinoline carboxylic acid. It is a slightly yellow to yellow crystalline substance with a molecular weight of 437.9. Its empirical formula is C₂₁H₂₄FN₃O₄ *HCl and its chemical structure is as follows:

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Moxifloxacin differs from other quinolones in that it has a methoxy function at the

8-position, and an S,S – configured diazabicyclononyl ring moiety at the 7-position.

AVELOX is available in 400 mg (moxifloxacin equivalent) film-coated tablets. The

inactive ingredients are microcrystalline cellulose, lactose monohydrate,

titanium dioxide, polyethylene glycol and ferric oxide.

croscarmellose sodium, magnesium stearate, hydroxypropyl methylcellulose,

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CLINICAL PHARMACOLOGY

Absorption

Moxifloxacin, given as an oral tablet, is well absorbed from the gastrointestinal 28 tract. The absolute bioavailability of moxifloxacin is approximately 90 percent. Co-administration with a high fat meal (i.e., 500 calories from fat) does not affect the absorption of moxifloxacin.

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Consumption of 1 cup of yogurt with moxifloxacin does not significantly affect the extent or rate of systemic absorption (AUC).

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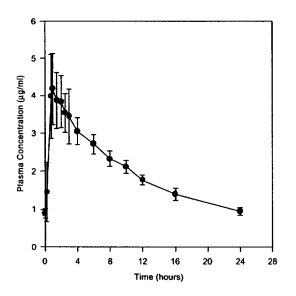
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The mean (± SD) C_{max} and AUC values at steady-state with a 400 mg once daily dosage regimen are $4.5 \pm 0.53 \,\mu\text{g/mL}$ and $48 \pm 2.7 \,\mu\text{g}^*\text{h/mL}$, respectively. C_{max} is attained 1 to 3 hours after oral dosing. The mean (± SD) trough concentration is 0.95 ± 0.10 μg/mL. Plasma concentrations increase proportionately with dose up

to the highest dose tested (800 mg single dose). The mean (\pm SD) elimination half-life from plasma is 12 \pm 1.3 hours; steady-state is achieved after at least three days with a 400 mg once daily regimen. The figure below illustrates the time course of plasma concentrations of moxifloxacin following a 400 mg dose administered at steady-state.

Steady-State Plasma Concentrations of Moxifloxacin Obtained With Once Daily Dosing of 400 mg (mean;SD) (n=10)



45 **Distribution**

Moxifloxacin is approximately 50% bound to serum proteins, independent of drug 46 concentration. The volume of distribution of moxifloxacin ranges from 1.7 to 2.7 47 L/kg. Moxifloxacin is widely distributed throughout the body, with tissue 48 concentrations often exceeding plasma concentrations. Moxifloxacin has been 49 detected in the saliva, nasal and bronchial secretions, mucosa of the sinuses, skin 50 blister fluid, and subcutaneous tissue, and skeletal muscle following oral 51 administration of 400 mg. Concentrations measured at 3 hours post-dose are 52 summarized in the following table. The rates of elimination of moxifloxacin from 53 54 tissues generally parallel the elimination from plasma.

Moxifloxacin Concentrations (mean \pm SD) in Plasma and Tissues Measured 3 Hours After Dosing with 400 mg §

Tissue or Fluid	N	Plasma Concentration (μg/mL)	Tissue or Fluid Concentration (μg/mL or μg/g)	Tissue: Plasma Ratio
Respiratory				
Alveolar Macrophages	5	3.3 ± 0.7	61.8 ± 27.3	21.2 ± 10.0
Bronchial Mucosa	8	3.3 ± 0.7	5.5 ± 1.3	1.7 ± 0.3
Epithelial Lining Fluid	5	3.3 ± 0.7	24.4 ± 14.7	8.7 ± 6.1
Sinus				
Maxillary Sinus Mucosa	4	3.7 ± 1.1 [†]	7.6 ± 1.7	2.0 ± 0.3
Anterior Ethmoid Mucosa	3	3.7 ± 1.1 [†]	8.8 ± 4.3	2.2 ± 0.6
Nasal Polyps	4	3.7 ± 1.1 [†]	9.8 ± 4.5	2.6 ± 0.6

 ^{\$} all moxifloxacin concentrations were measured after a single 400 mg dose,
 except the sinus concentrations which were measured after 5 days of dosing.

Metabolism

Moxifloxacin is metabolized via glucuronide and sulfate conjugation. The cytochrome P450 system is not involved in moxifloxacin metabolism, and is not affected by moxifloxacin. The sulfate conjugate (M1) accounts for approximately 38% of the dose, and is eliminated primarily in the feces. Approximately 14% of an oral or intravenous dose is converted to a glucuronide conjugate (M2), which is excreted exclusively in the urine. Peak plasma concentrations of M2 are approximately 40% those of the parent drug, while plasma concentrations of M1 are generally less than 10% those of moxifloxacin.

Excretion

Approximately 45% of an oral or intravenous dose of moxifloxacin is excreted as unchanged drug (~20% in urine and ~25% in feces). A total of 96% \pm 4% of an oral dose is excreted as either unchanged drug or known metabolites. The mean (\pm SD) apparent total body clearance and renal clearance are 12 \pm 2.0 L/hr and 2.6 \pm 0.5 L/hr, respectively.

Special Populations

Geriatric

In 16 healthy elderly male and female volunteers (66-81 years of age) given a single 200 mg dose of moxifloxacin, the extent of systemic exposure (AUC and C_{max}) was not statistically different between young and elderly males and elimination half-life was unchanged. No dosage adjustment is necessary based on age.

[†] N = 5

Whether pharmacokinetic differences exist between young and elderly females is unknown. The pharmacokinetics of moxifloxacin with repeated 400 mg administration in elderly subjects has not been studied.

Pediatric

The pharmacokinetics of moxifloxacin in pediatric subjects have not been studied.

Gender

Following a single 200 mg dose of moxifloxacin to 16 healthy elderly subjects, the mean AUC and C_{max} were 29% and 24% higher, respectively, in healthy elderly females compared to healthy elderly males. There are no significant differences in moxifloxacin pharmacokinetics between elderly male and female subjects when differences in body weight are taken into consideration.

A 400 mg single dose study was conducted in 18 young males and females. The comparison of moxifloxacin pharmacokinetics in this study (9 young females and 9 young males) showed no differences in AUC or Cmax due to gender. Dosage adjustments based on gender are not necessary.

Race

Steady state moxifloxacin pharmacokinetics in male Japanese subjects were similar to those determined in Caucasians, with a mean C_{max} of 4.1 μ g/mL, an AUC₂₄ of 47 μ g*h/mL, and an elimination half-life of 14 hours.

Renal Insufficiency

The pharmacokinetic parameters of moxifloxacin are not significantly altered by mild, moderate, or severe renal impairment. No dosage adjustment is necessary in patients with renal impairment.

 In a single-dose study of 24 patients with varying degrees of renal function from normal to severely impaired, the mean peak concentrations (C_{max}) of moxifloxacin were reduced by 22% and 21% in the patients with moderate ($CL_{CR} \geq 30$ and ≤ 60 mL/min) and severe ($CL_{CR} < 30$ mL/min) renal impairment, respectively. The mean systemic exposure (AUC) in these patients was increased by 13%. In the moderate and severe renally impaired patients, the mean AUC for the sulfate conjugate (M1) increased by 1.7-fold (ranging up to 2.8-fold) and mean AUC and Cmax for the glucuronide conjugate (M2) increased by 2.8-fold (ranging up to 4.8-fold) and 1.4-fold (ranging up to 2.5-fold), respectively. The sulfate and glucuronide conjugates are not microbiologically active, and the clinical implication of increased exposure to these metabolites in patients with renal impairment has not been studied.

The effect of hemodialysis or continuous ambulatory peritoneal dialysis (CAPD) on the pharmacokinetics of moxifloxacin has not been studied.

Hepatic Insufficiency

In 400 mg single dose studies in 6 patients with mild (Child Pugh Class A), and 10 patients with moderate (Child Pugh Class B), hepatic insufficiency, moxifloxacin mean systemic exposure (AUC) was 78% and 102%, respectively, of 18 healthy controls and mean peak concentration (C_{max}) was 79% and 84% of controls.

The mean AUC of the sulfate conjugate of moxifloxacin (M1) increased by 3.9-fold (ranging up to 5.9-fold) and 5.7-fold (ranging up to 8.0-fold) in the mild and moderate groups, respectively. The mean C_{max} of M1 increased by approximately 3-fold in both groups (ranging up to 4.7- and 3.9-fold). The mean AUC of the glucuronide conjugate of moxifloxacin (M2) increased by 1.5-fold (ranging up to 2.5-fold) in both groups. The mean C_{max} of M2 increased by 1.6-and 1.3-fold (ranging up to 2.7- and 2.1-fold), respectively. The clinical significance of increased exposure to the sulfate and glucuronide conjugates has not been studied. No dosage adjustment is recommended for mild or moderate hepatic insufficiency (Child Pugh Classes A and B). The pharmacokinetics of moxifloxacin in severe hepatic insufficiency (Child Pugh Class C) have not been studied. (See **DOSAGE AND ADMINISTRATION**.)

Photosensitivity Potential

A study of the skin response to ultraviolet (UVA and UVB) and visible radiation conducted in 32 healthy volunteers (8 per group) demonstrated that moxifloxacin does not show phototoxicity in comparison to placebo. The minimum erythematous dose (MED) was measured before and after treatment with moxifloxacin (200 mg or 400 mg once daily), lomefloxacin (400 mg once daily), or placebo. In this study, the MED measured for both doses of moxifloxacin were not significantly different from placebo, while lomefloxacin significantly lowered the MED. (See **PRECAUTIONS**, **Information for Patients**.)

Drug-drug interactions

The potential for pharmacokinetic drug interactions between moxifloxacin and theophylline, warfarin, digoxin, probenecid, ranitidine, glyburide, iron, and antacids has been evaluated. There was no clinically significant effect of moxifloxacin on theophylline, warfarin, digoxin, or glyburide kinetics. Theophylline, digoxin, probenecid, and ranitidine did not affect the pharmacokinetics of moxifloxacin. However, as with all other quinolones, iron and antacids significantly reduced the bioavailability of moxifloxacin.

Theophylline: No significant effect of moxifloxacin (200 mg every twelve hours for 3 days) on the pharmacokinetics of theophylline (400 mg every twelve hours for 3 days) was detected in a study involving 12 healthy volunteers. In addition, theophylline was not shown to affect the pharmacokinetics of moxifloxacin. The effect of co-administration of a 400 mg dose of moxifloxacin with theophylline has not been studied, but it is not expected to be clinically significant based on *in vitro* metabolic data showing that moxifloxacin does not inhibit the CYP1A2 isoenzyme.

Warfarin: No significant effect of moxifloxacin (400 mg once daily for eight days) on the pharmacokinetics of R- and S-warfarin (25 mg single dose of warfarin sodium on the fifth day) was detected in a study involving 24 healthy volunteers. No significant change in prothrombin time was observed. (See PRECAUTIONS, Drug Interactions.)

 Digoxin: No significant effect of moxifloxacin (400 mg once daily for two days) on digoxin (0.6 mg as a single dose) AUC was detected in a study involving 12 healthy volunteers. The mean digoxin C_{max} increased by about 50% during the distribution phase of digoxin. This transient increase in digoxin C_{max} is not viewed to be clinically significant. Moxifloxacin pharmacokinetics were similar in the presence or absence of digoxin. No dosage adjustment for moxifloxacin or digoxin is required when these drugs are administered concomitantly.

Probenecid: Probenecid (500 mg twice daily for two days) did not alter the renal clearance and total amount of moxifloxacin (400 mg single dose) excreted renally in a study of 12 healthy volunteers.

Ranitidine: No significant effect of ranitidine (150 mg twice daily for three days as pretreatment) on the pharmacokinetics of moxifloxacin (400 mg single dose) was detected in a study involving 10 healthy volunteers.

Antidiabetic agents: In diabetics, glyburide (2.5 mg once daily for two weeks pretreatment and for five days concurrently) mean AUC and C_{max} were 12% and 21 % lower, respectively, when taken with moxifloxacin (400 mg once daily for five days) in comparison to placebo. Nonetheless, blood glucose levels were decreased slightly in patients taking glyburide and moxifloxacin in comparison to those taking glyburide alone, suggesting no interference by moxifloxacin on the activity of glyburide. These interaction results are not viewed as clinically significant.

Antacids: When moxifloxacin (single 400 mg dose) was administered two hours before, concomitantly, or 4 hours after an aluminum/magnesium-containing antacid (900 mg aluminum hydroxide and 600 mg magnesium hydroxide as a single oral dose) to 12 healthy volunteers there was a 26%, 60% and 23% reduction in the mean AUC of moxifloxacin, respectively. Moxifloxacin should be taken at least 4 hours before or 8 hours after antacids containing magnesium or aluminum, as well as sucralfate, metal cations such as iron, and multivitamin preparations with zinc, or Videx® (didanosine) chewable/buffered tablets or the pediatric powder for oral solution. (See PRECAUTIONS, Drug Interactions and DOSAGE AND ADMINISTRATION.)

Iron: When moxifloxacin was administered concomitantly with iron (ferrous sulfate 100 mg once daily for two days), the mean AUC and C_{max} of moxifloxacin was reduced by 39% and 59%, respectively. Moxifloxacin should only be taken more

225 226	than 4 hours before or 8 hours after iron products. (See PRECAUTIONS, Drug Interactions and DOSAGE AND ADMINISTRATION.)
227	
228	There is limited information available on the potential for a pharmacodynamic
229	interaction in humans between moxifloxacin and other drugs that prolong the QTc
230	interval of the electrocardiogram. Sotalol, a Class III antiarrhythmic, has been
231	shown to further increase the QTc interval when combined with high doses of
232	intravenous (IV) moxifloxacin in dogs. Therefore, moxifloxacin should be avoided
233	with Class IA and Class III antiarrhythmics. (See ANIMAL PHARMACOLOGY,
234	WARNINGS, and PRECAUTIONS.)
235	
236	MICROBIOLOGY
237	Moxifloxacin has in vitro activity against a wide range of Gram-positive and Gram-
238	negative microorganisms. The bactericidal action of moxifloxacin results from
239	inhibition of the topoisomerase II (DNA gyrase) and topoisomerase IV required for
240	bacterial DNA replication, transcription, repair, and recombination. It appears
241	that the C8-methoxy moiety contributes to enhanced activity and lower selection of
242	resistant mutants of Gram-positive bacteria compared to the C8-H moiety.
243	,
244	The mechanism of action for quinolones, including moxifloxacin, is different from
245	that of macrolides, beta-lactams, aminoglycosides, or tetracyclines; therefore,
246	microorganisms resistant to these classes of drugs may be susceptible to
247	moxifloxacin and other quinolones. There is no known cross-resistance between
248	moxifloxacin and other classes of antimicrobials.
249	
250	Cross-resistance has been observed between moxifloxacin and other
251	fluoroquinolones against Gram-negative bacteria. Gram-positive bacteria
252	resistant to other fluoroquinolones may, however, still be susceptible to
253	moxifloxacin.
254	
255	Moxifloxacin has been shown to be active against most strains of the following
256	microorganisms, both in vitro and in clinical infections as described in the
257	INDICATIONS AND USAGE section.
258	
259	Aerobic Gram-positive microorganisms
260	Staphylococcus aureus (methicillin-susceptible strains only)
261	Streptococcus pneumoniae (penicillin-susceptible strains)
262	Streptococcus pyogenes
263	, , , , , , , , , , , , , , , , , , , ,
264	Aerobic Gram-negative microorganisms
265	Haemophilus influenzae
266	Haemophilus parainfluenzae
267	Klebsiella pneumoniae
268	Moraxella catarrhalis
269	

Other microorganisms

271	Chlamydia pneumoniae
272	Mycoplasma pneumoniae
273	
274	The following in vitro data are available, but their clinical significance is
275	<u>unknown</u> .
276	
277	Moxifloxacin exhibits in vitro minimum inhibitory concentrations (MICs) of 2
278	μg/mL or less against most (≥90%) strains of the following microorganisms;
279	however, the safety and effectiveness of moxifloxacin in treating clinical infections
280	due to these microorganisms have not been established in adequate and well-
281	controlled clinical trials.
282	
283	Aerobic Gram-positive microorganisms
284	Staphylococcus epidermidis (methicillin-susceptible strains only)
285	Streptococcus agalactiae
286	Streptococcus pneumoniae (penicillin-resistant strains)
287	Streptococcus viridans group
288	
289	Aerobic Gram-negative microorganisms
290	Citrobacter freundii
291	Enterobacter cloacae
292	Escherichia coli
293	Klebsiella oxytoca
294	Legionella pneumophila
295	Proteus mirabilis
296	
297	Anaerobic microorganisms
298	Fusobacterium species
299	Peptostreptococcus species
300	Prevotella species
301	
302	Susceptibility Tests
303	Dilution Techniques : Quantitative methods are used to determine antimicrobial
304	minimum inhibitory concentrations (MICs). These MICs provide estimates of the
305	susceptibility of bacteria to antimicrobial compounds. The MICs should be
306	determined using a standardized procedure. Standardized procedures are based
307	on a dilution method¹ (broth or agar) or equivalent with standardized inoculum
308	concentrations and standardized concentrations of moxifloxacin powder. The MIC

values should be interpreted according to the following criteria:

For testing Enterobacteriaceae and *Staphylococcus* species:

313	MIC (μg/mL)	<u>Interpretation</u>
314	≤ 2.0	Susceptible (S)
315	4.0	Intermediate (I)
316	≥ 8.0	Resistant (R)

For testing Haemophilus influenzae and Haemophilus parainfluenzae a:

320	MIC (μg/mL)	<u>Interpretation</u>
321	<u>≤ 1.0</u>	Susceptible (S)

^a This interpretive standard is applicable only to broth microdilution susceptibility tests with *Haemophilus influenzae* and *Haemophilus parainfluenzae* using *Haemophilus* Test Medium¹.

The current absence of data on resistant strains precludes defining any results other than "Susceptible". Strains yielding MIC results suggestive of a "nonsusceptible " category should be submitted to a reference laboratory for further testing.

For testing Streptococcus species including Streptococcus pneumoniae b:

334	<u>MIC (μg/mL)</u>	<u>Interpretation</u>
335	≤ 1.0	Susceptible (S)
336	2.0	Intermediate (I)
337	≥ 4.0	Resistant (R)

^b This interpretive standard is applicable only to broth microdilution susceptibility tests using cation-adjusted Mueller-Hinton broth with 2 - 5% lysed horse blood.

A report of "Susceptible" indicates that the pathogen is likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable. A report of "Intermediate" indicates that the result should be considered equivocal, and, if the microorganism is not fully susceptible to alternative, clinically feasible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is physiologically concentrated or in situations where a high dosage of drug can be used. This category also provides a buffer zone which prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of "Resistant" indicates that the pathogen is not likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable; other therapy should be selected.

Standardized susceptibility test procedures require the use of laboratory control microorganisms to control the technical aspects of the laboratory procedures.

Standard moxifloxacin powder should provide the following MIC values:

359	<u>Microorganism</u>		MIC (µg/mL)
360	Enterococcus faecalis	ATCC 29212	0.06 - 0.5
361	Escherichia coli	ATCC 25922	0.008 - 0.06
362	Haemophilus influenzae	ATCC 49247°	0.008 - 0.03
363	Staphylococcus aureus	ATCC 29213	0.015 - 0.06
364	Streptococcus pneumoniae	ATCC 49619 ^d	0.06 - 0.25

^c This quality control range is applicable to only *H. influenzae* ATCC 49247 tested by a broth microdilution procedure using *Haemophilus* Test Medium (HTM)¹.

^d This quality control range is applicable to only *S. pneumoniae* ATCC 49619 tested by a broth microdilution procedure using cation-adjusted Mueller-Hinton broth with 2 - 5% lysed horse blood.

Diffusion Techniques: Quantitative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of bacteria to antimicrobial compounds. One such standardized procedure² requires the use of standardized inoculum concentrations. This procedure uses paper disks impregnated with 5-µg moxifloxacin to test the susceptibility of microorganisms to moxifloxacin.

Reports from the laboratory providing results of the standard single-disk susceptibility test with a 5-µg moxifloxacin disk should be interpreted according to the following criteria:

The following zone diameter interpretive criteria should be used for testing Enterobacteriaceae and *Staphylococcus* species:

Zone Diameter (mm)	<u>Interpretation</u>
≥ 19	Susceptible (S)
16 - 18	Intermediate (I)
≤ 15	Resistant (R)

For testing Haemophilus influenzae and Haemophilus parainfluenzae e:

Zone Diameter (mm)	<u>Interpretation</u>
≥ 18	Susceptible (S)

^eThis zone diameter standard is applicable only to tests *with Haemophilus influenzae* and *Haemophilus parainfluenzae* using *Haemophilus* Test Medium (HTM)².

The current absence of data on resistant strains precludes defining any results other than "Susceptible". Strains yielding zone diameter results suggestive of a "nonsusceptible" category should be submitted to a reference laboratory for further testing.

For testing Streptococcus species including Streptococcus pneumoniae^f:

409	Zone Diameter (mm)	<u>Interpretation</u>
410	≥ 18	Susceptible (S)
411	15 - 17	Intermediate (I)
412	≤ 14	Resistant (R)

^f These interpretive standards are applicable only to disk diffusion tests using Mueller-Hinton agar supplemented with 5% sheep blood incubated in 5% CO₂.

Interpretation should be as stated above for results using dilution techniques. Interpretation involves correlation of the diameter obtained in the disk test with the MIC for moxifloxacin.

As with standardized dilution techniques, diffusion methods require the use of laboratory control microorganisms that are used to control the technical aspects of the laboratory procedures. For the diffusion technique, the 5-µg moxifloxacin disk should provide the following zone diameters in these laboratory test quality control strains:

427	<u>Microorganism</u>		Zone Diameter (mm)
428	Escherichia coli	ATCC 25922	28 - 35
429	Haemophilus influenzae	ATCC 49274 ⁹	31 - 39
430	Staphylococcus aureus	ATCC 25923	28 - 35
431	Streptococcus pneumoniae	ATCC 49619 ^h	25 - 31

⁹These quality control limits are applicable to only *H. influenzae* ATCC 49247 testing using *Haemophilus* Test Medium (HTM)².

^hThese quality control limits are applicable only to tests conducted with *S. pneumoniae* ATCC 49619 performed by disk diffusion using Mueller-Hinton agar supplemented with 5% defibrinated sheep blood.

INDICATIONS AND USAGE

AVELOX Tablets are indicated for the treatment of adults (≥ 18 years of age) with infections caused by susceptible strains of the designated microorganisms in the

443	conditions listed below. Please see DOSAGE AND ADMINISTRATION for
444	specific recommendations.
445	
446	Acute Bacterial Sinusitis caused by Streptococcus pneumoniae, Haemophilus
447	influenzae, or Moraxella catarrhalis.
448	
449	Acute Bacterial Exacerbation of Chronic Bronchitis caused by Streptococcus
450	pneumoniae, Haemophilus influenzae, Haemophilus parainfluenzae, Klebsiella
451	pneumoniae, Staphylococcus aureus, or Moraxella catarrhalis.
452	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
453	Community Acquired Pneumonia (of mild to moderate severity) caused by
454	Streptococcus pneumoniae, Haemophilus influenzae, Mycoplasma pneumoniae,
455	Chlamydia pneumoniae, or Moraxella catarrhalis.
456	
457	Uncomplicated Skin and Skin Structure Infections caused by Staphylococcus
458	aureus or Streptococcus pyogenes.
459	, , , , ,
460	Appropriate culture and susceptibility tests should be performed before treatment
461	in order to isolate and identify organisms causing infection and to determine their
462	susceptibility to moxifloxacin. Therapy with AVELOX may be initiated before
463	results of these tests are known; once results become available, appropriate
464	therapy should be continued.
465	
466	CONTRAINDICATIONS
467	
468	Moxifloxacin is contraindicated in persons with a history of hypersensitivity to
469	moxifloxacin or any member of the quinolone class of antimicrobial agents.
470	
471	WARNINGS
472	
473	THE SAFETY AND EFFECTIVENESS OF MOXIFLOXACIN IN PEDIATRIC
474	PATIENTS, ADOLESCENTS (LESS THAN 18 YEARS OF AGE), PREGNANT
475	WOMEN, AND LACTATING WOMEN HAVE NOT BEEN ESTABLISHED. (SEE
476	PRECAUTIONS-PEDIATRIC USE, PREGNANCY AND NURSING MOTHERS
477	SUBSECTIONS.)
478	
479	MOXIFLOXACIN HAS BEEN SHOWN TO PROLONG THE QT INTERVAL OF
480	THE ELECTROCARDIOGRAM IN SOME PATIENTS. THE DRUG SHOULD BE
481	AVOIDED IN PATIENTS WITH KNOWN PROLONGATION OF THE QT
482	INTERVAL, PATIENTS WITH UNCORRECTED HYPOKALEMIA AND
483	PATIENTS RECEIVING CLASS IA (E.G. QUINIDINE, PROCAINAMIDE) OR
484	CLASS III (E.G. AMIODARONE, SOTALOL) ANTIARRHYTHMIC AGENTS, DUE
485	TO THE LACK OF CLINICAL EXPERIENCE WITH THE DRUG IN THESE
486	PATIENT POPULATIONS.
487	

Pharmacokinetic studies between moxifloxacin and other drugs that prolong the QT interval such as cisapride, erythromycin, antipsychotics, and tricyclic antidepressants have not been performed. An additive effect of moxifloxacin and these drugs cannot be excluded, therefore moxifloxacin should be used with caution when given concurrently with these drugs.

The effect of moxifloxacin on patients with congenital prolongation of the QT interval has not been studied, however, it is expected that these individuals may be more susceptible to drug-induced QT prolongation. Because of limited clinical experience, moxifloxacin should be used with caution in patients with ongoing proarrhythmic conditions, such as clinically significant bradycardia, acute myocardial ischemia.

The magnitude of QT prolongation may increase with increasing concentrations of the drug, therefore the recommended dose should not be exceeded. QT prolongation may lead to an increased risk for ventricular arrhythmias including torsade de pointes. In 787 patients with paired valid ECGs in Phase III clinical trials, the mean \pm SD effect of moxifloxacin 400 mg on the QTc interval was 6 \pm 26 msec. No cardiovascular morbidity or mortality attributable to QTc prolongation occurred with moxifloxacin treatment in over 4000 patients in controlled clinical studies, and there was no increase in mortality in over 18,000 patients in a post-marketing observational study. However certain predisposing conditions may increase the risk for ventricular arrhythmias.

The oral administration of moxifloxacin caused lameness in immature dogs. Histopathological examination of the weight-bearing joints of these dogs revealed permanent lesions of the cartilage. Related quinolone-class drugs also produce erosions of cartilage of weight-bearing joints and other signs of arthropathy in immature animals of various species. (See **ANIMAL PHARMACOLOGY**.)

Convulsions have been reported in patients receiving quinolones. Quinolones may also cause central nervous system (CNS) events including: dizziness, confusion, tremors, hallucinations, depression, and, rarely, suicidal thoughts or acts. These reactions may occur following the first dose. If these reactions occur in patients receiving moxifloxacin, the drug should be discontinued and appropriate measures instituted. As with all quinolones, moxifloxacin should be used with caution in patients with known or suspected CNS disorders (e.g. severe cerebral arteriosclerosis, epilepsy) or in the presence of other risk factors that may predispose to seizures or lower the seizure threshold. (See PRECAUTIONS: General, Information for Patients, and ADVERSE REACTIONS.)

Serious anaphylactic reactions, some following the first dose, have been reported in patients receiving quinolone therapy, including moxifloxacin. Some reactions were accompanied by cardiovascular collapse, loss of consciousness, tingling, pharyngeal or facial edema, dyspnea, urticaria, and itching. Serious anaphylactic reactions require immediate emergency treatment with epinephrine. Moxifloxacin

should be discontinued at the first appearance of a skin rash or any other sign of hypersensitivity. Oxygen, intravenous steroids, and airway management, including intubation, may be administered as indicated.

Severe and sometimes fatal events, some due to hypersensitivity, and some of uncertain etiology, have been reported in patients receiving therapy with all antibiotics. These events may be severe and generally occur following the administration of multiple doses. Clinical manifestations may include one or more of the following: rash, fever, eosinophilia, jaundice, and hepatic necrosis.

Pseudomembranous colitis has been reported with nearly all antibacterial agents and may range in severity from mild to life-threatening. Therefore, it is important to consider this diagnosis in patients who present with diarrhea subsequent to the administration of antibacterial agents.

Treatment with antibacterial agents alters the normal flora of the colon and may permit overgrowth of clostridia. Studies indicate that a toxin produced by *Clostridium difficile* is one primary cause of "antibiotic-associated colitis."

After the diagnosis of pseudomembranous colitis has been established, therapeutic measures should be initiated. Mild cases of pseudomembranous colitis usually respond to drug discontinuation alone. In moderate to severe cases, consideration should be given to management with fluids and electrolytes, protein supplementation, and treatment with an antibacterial drug clinically effective against *C. difficile* colitis.

Although not observed in moxifloxacin clinical trials, Achilles and other tendon ruptures that required surgical repair or resulted in prolonged disability have been reported with quinolones. Moxifloxacin should be discontinued if the patient experiences pain, inflammation, or rupture of a tendon.

PRECAUTIONS

General: Quinolones may cause central nervous system (CNS) events, including: nervousness, agitation, insomnia, anxiety, nightmares or paranoia. (See **WARNINGS** and **Information for Patients**.)

Information for Patients:

To assure safe and effective use of moxifloxacin, the following information and instructions should be communicated to the patient when appropriate:

Patients should be advised:

 that moxifloxacin may produce changes in the electrocardiogram (QTc interval prolongation).

that moxifloxacin should be avoided in patients receiving Class IA (e.g.
 quinidine, procainamide) or Class III (e.g. amiodarone, sotalol) antiarrhythmic
 agents.

 that moxifloxacin may add to the QTc prolonging effects of other drugs such as cisapride, erythromycin, antipsychotics, and tricyclic antidepressants.

• to inform their physician of any personal or family history of QTc prolongation or proarrhythmic conditions such as recent hypokalemia, significant bradycardia, acute myocardial ischemia.

• to inform their physician of any other medications when taken concurrently with moxifloxacin, including over-the-counter medications.

• to contact their physician if they experience palpitations or fainting spells while taking moxifloxacin.

 that moxifloxacin may be taken with or without meals, and to drink fluids liberally.

that moxifloxacin should be taken at least 4 hours before or 8 hours after multivitamins (containing iron or zinc), antacids (containing magnesium, calcium, or aluminum), sucralfate, or Videx® (didanosine) chewable/buffered tablets or the pediatric powder for oral solution. (See CLINICAL PHARMACOLOGY, Drug Interactions and PRECAUTIONS, Drug Interactions.)

 that moxifloxacin may be associated with hypersensitivity reactions, including anaphylactic reactions, even following a single dose, and to discontinue the drug at the first sign of a skin rash or other signs of an allergic reaction.

◆ to discontinue treatment; rest and refrain from exercise; and inform their physician if they experience pain, inflammation, or rupture of a tendon.

 that moxifloxacin may cause dizziness and lightheadedness; therefore, patients should know how they react to this drug before they operate an automobile or machinery or engage in activities requiring mental alertness or coordination.

 that phototoxicity has been reported in patients receiving certain quinolones. There was no phototoxicity seen with moxifloxacin at the recommended dose. In keeping with good medical practice, avoid excessive sunlight or artificial ultraviolet light (e.g. tanning beds). If sunburn-like reaction or skin eruptions occur, contact your physician. (See CLINICAL PHARMACOLOGY, Photosensitivity Potential.)

 that convulsions have been reported in patients receiving quinolones, and they should notify their physician before taking this drug if there is a history of this condition.

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Drug Interactions:

- 628 Antacids, Sucralfate, Metal Cations, Multivitamins: Quinolones form chelates with
- alkaline earth and transition metal cations. Administration of quinolones with
- antacids containing aluminum, magnesium, or calcium, with sucralfate, with metal
- cations such as iron, or with multivitamins containing iron or zinc, or with
- formulations containing divalent and trivalent cations such as Videx[®] (didanosine)
- chewable/buffered tablets or the pediatric powder for oral solution, may
- substantially interfere with the absorption of guinolones, resulting in systemic
- 635 concentrations considerably lower than desired. Therefore, moxifloxacin should
- be taken at least 4 hours before or 8 hours after these agents. (See CLINICAL
 - PHARMACOLOGY, Drug Interactions and DOSAGE AND ADMINISTRATION.)

No clinically significant drug-drug interactions between theophylline, warfarin, digoxin, or glyburide have been observed with moxifloxacin. Theophylline, digoxin, probenecid, and ranitidine have been shown not to alter the pharmacokinetics of moxifloxacin. (See CLINICAL PHARMACOLOGY.)

Warfarin: No significant effect of moxifloxacin on R- and S- warfarin was detected in a clinical study involving 24 healthy volunteers. No significant changes in prothrombin time were noted in the presence of moxifloxacin. However, since some quinolones have been reported to enhance the anticoagulant effects of warfarin or its derivatives in the patient population, the prothrombin time or other suitable coagulation test should be closely monitored if a quinolone antimicrobial is administered concomitantly with warfarin or its derivatives.

Drugs metabolized by Cytochrome P450 enzymes: *In vitro* studies with cytochrome P450 isoenzymes (CYP) indicate that moxifloxacin does not inhibit CYP3A4, CYP2D6, CYP2C9, CYP2C19, or CYP1A2, suggesting that moxifloxacin is unlikely to alter the pharmacokinetics of drugs metabolized by

these enzymes (e.g. midazolam, cyclosporine, warfarin, theophylline).

Nonsteroidal anti-inflammatory drugs (NSAIDs): Although not observed with moxifloxacin in preclinical and clinical trials, the concomitant administration of a nonsteroidal anti-inflammatory drug with a quinolone may increase the risks of CNS stimulation and convulsions. (See **WARNINGS**.)

Carcinogenesis, Mutagenesis, Impairment of Fertility:

Long term studies in animals to determine the carcinogenic potential of moxifloxacin have not been performed.

Moxifloxacin was not mutagenic in 4 bacterial strains (TA 98, TA 100, TA 1535, TA 1537) used in the Ames *Salmonella* reversion assay. As with other quinolones, the positive response observed with moxifloxacin in strain TA 102 using the same assay may be due to the inhibition of DNA gyrase. Moxifloxacin was not mutagenic in the CHO/HGPRT mammalian cell gene mutation assay. An equivocal result was obtained in the same assay when v79 cells were used. Moxifloxacin was

clastogenic in the v79 chromosome aberration assay, but it did not induce unscheduled DNA synthesis in cultured rat hepatocytes. There was no evidence of genotoxicity *in vivo* in a micronucleus test or a dominant lethal test in mice.

Moxifloxacin had no effect on fertility in male and female rats at oral doses as high as 500 mg/kg/day, approximately 12 times the maximum recommended human dose based on body surface area (mg/m²). At 500 mg/kg there were slight effects on sperm morphology (head-tail separation) in male rats and on the estrous cycle in female rats.

Pregnancy: Teratogenic Effects. Pregnancy Category C:

Moxifloxacin was not teratogenic when administered to pregnant rats during organogenesis at oral doses as high as 500 mg/kg/day or 0.24 times the maximum recommended human dose based on systemic exposure (AUC), but decreased fetal body weights and slightly delayed fetal skeletal development (indicative of fetotoxicity) were observed. Intravenous administration of 20 mg/kg/day (approximately equal to the maximum recommended human oral dose based upon systemic exposure) to pregnant rabbits during organogenesis resulted in decreased fetal body weights and delayed fetal skeletal ossification. When rib and vertebral malformations were combined, there was an increased fetal and litter incidence of these effects. Signs of maternal toxicity in rabbits at this dose included mortality, abortions, marked reduction of food consumption, decreased water intake, body weight loss and hypoactivity. There was no evidence of teratogenicity when pregnant Cynomolgus monkeys were given oral doses as high as 100 mg/kg/day (2.5 times the maximum recommended human dose based upon systemic exposure). An increased incidence of smaller fetuses was observed at 100 mg/kg/day. In an oral pre- and postnatal development study conducted in rats, effects observed at 500 mg/kg/day included slight increases in duration of pregnancy and prenatal loss, reduced pup birth weight and decreased neonatal survival. Treatment-related maternal mortality occurred during gestation at 500 mg/kg/day in this study.

Since there are no adequate or well-controlled studies in pregnant women, moxifloxacin should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Nursing Mothers: Moxifloxacin is excreted in the breast milk of rats. Moxifloxacin may also be excreted in human milk. Because of the potential for serious adverse reactions in infants nursing from mothers taking moxifloxacin, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

Pediatric Use: Safety and effectiveness in pediatric patients and adolescents less than 18 years of age have not been established. Moxifloxacin causes arthropathy in juvenile animals. (See **WARNINGS**.)

718 Geriatric Use: In controlled multiple-dose clinical trials, 23% of patients receiving moxifloxacin were greater than or equal to 65 years of age and 9% were greater 719 than or equal to 75 years of age. The clinical trial data demonstrate that there is 720 no difference in the safety and efficacy of moxifloxacin in patients aged 65 or older 721 compared to younger adults. 722

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ADVERSE REACTIONS

Clinical efficacy trials enrolled over 4900 moxifloxacin treated patients, of whom over 4300 patients received the 400 mg dose. Most adverse events reported in moxifloxacin trials were described as mild to moderate in severity and required no treatment. Moxifloxacin was discontinued due to adverse reactions thought to be drug-related in 3.8% of patients.

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Adverse reactions, judged by investigators to be at least possibly drug-related, occurring in greater than or equal to 1% of moxifloxacin treated patients were: 732 nausea (8%), diarrhea (6%), dizziness (3%), headache (2%), abdominal pain (2%), vomiting (2%), taste perversion (1%), abnormal liver function test (1%), and dyspepsia (1%).

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Additional events, judged by investigators to be at least possibly drug-related, that occurred in greater than 0.05% and less than 1% of moxifloxacin treated patients were:

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- 741 BODY AS A WHOLE: asthenia, moniliasis, pain, malaise, lab test abnormal (not 742 specified), allergic reaction, leg pain, pelvic pain, abdominal pain, back pain, chills, 743 infection, chest pain, hand pain
- CARDIOVASCULAR: palpitation, vasodilatation, tachycardia, hypertension, 744
- 745 peripheral edema, hypotension
- 746 CENTRAL NERVOUS SYSTEM: insomnia, nervousness, anxiety, confusion.
- hallucinations, depersonalization, hypertonia, incoordination, somnolence, tremor, 747
- 748 vertigo, paresthesia
- DIGESTIVE: dry mouth, constipation, oral moniliasis, anorexia, stomatitis, 749
- 750 gastritis, glossitis, gastrointestinal disorder, cholestatic jaundice, GGTP increased
- 751 HEMIC AND LYMPHATIC: prothrombin time decrease, prothrombin time increase.
- thrombocythemia, thrombocytopenia, eosinophilia, leukopenia 752
- 753 METABOLIC AND NUTRITIONAL: amylase increased, hyperglycemia.
- 754 hyperlipidemia, lactic dehydrogenase increased
- MUSCULOSKELETAL: arthralgia, myalgia 755
- 756 RESPIRATORY: asthma, dyspnea, cough increased, pneumonia, pharyngitis.
- rhinitis, sinusitis 757
- 758 SKIN/APPENDAGES: rash, pruritus, sweating, urticaria, dry skin.
- 759 SPECIAL SENSES: tinnitus, amblyopia
- 760 UROGENITAL: vaginal moniliasis, vaginitis, cystitis, kidney function abnormal

Post-Marketing Adverse Event Reports:

Additional adverse events reported from worldwide post-marketing experience with moxifloxacin include anaphylactic reaction and anaphylactic shock.

LABORATORY CHANGES

Changes in laboratory parameters, without regard to drug relationship, which are not listed above and which occurred in $\geq 2\%$ of patients and at an incidence greater than in controls included: increases in MCH, neutrophils, WBCs, PT ratio, ionized calcium, chloride, albumin, globulin, bilirubin; decreases in hemoglobin, RBCs, neutrophils, eosinophils, basophils, PT ratio, glucose, pO₂, bilirubin and amylase. It cannot be determined if any of the above laboratory abnormalities were caused by the drug or the underlying condition being treated.

OVERDOSAGE

In the event of acute overdosage, the stomach should be emptied and ECG monitoring is recommended due to the possible prolongation of the QT interval. The patient should be carefully observed and given supportive treatment. Adequate hydration must be maintained. It is not known whether moxifloxacin is dialyzable.

Single oral moxifloxacin doses of 2000, 500, and 1500 mg/kg were lethal to rats, mice, and cynomolgus monkeys, respectively. The minimum lethal intravenous dose in mice and rats was 100 mg/kg. Toxic signs after administration of a single high dose of moxifloxacin to these animals included CNS and gastrointestinal effects such as decreased activity, somnolence, tremor, convulsions, vomiting and diarrhea.

DOSAGE AND ADMINISTRATION

The dose of AVELOX Tablets is one 400 mg tablet taken orally every 24 hours. The duration of therapy depends on the type of infection as described below.

Infection *	Daily Dose	Duration
Acute Bacterial Sinusitis	400 mg	10 days
Acute Bacterial Exacerbation of Chronic Bronchitis	400 mg	5 days
Community Acquired Pneumonia	400 mg	10 days
Uncomplicated Skin and Skin Structure Infections	400 mg	7 days

^{*} due to the designated pathogens (See INDICATIONS AND USAGE.)

- 808 Oral doses of moxifloxacin should be administered at least 4 hours before or 8
- hours after antacids containing magnesium or aluminum, as well as sucralfate,
- metal cations such as iron, and multivitamin preparations with zinc, or Videx®
- (didanosine) chewable/buffered tablets or the pediatric powder for oral solution.
- (See CLINICAL PHARMACOLOGY, Drug Interactions and PRECAUTIONS,
- 813 **Drug Interactions.**)

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Impaired Renal Function

- No dosage adjustment is required in renally impaired patients. Moxifloxacin has not been studied in patients on hemodialysis or continuous ambulatory peritoneal
- 818 dialysis (CAPD).

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Impaired Hepatic Function

- No dosage adjustment is required in patients with mild or moderate hepatic
- insufficiency (Child Pugh Classes A and B). The pharmacokinetics of moxifloxacin
- in patients with severe hepatic insufficiency (Child Pugh Class C) have not been
- studied. (See CLINICAL PHARMACOLOGY, Hepatic Insufficiency.)

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HOW SUPPLIED

- 827 AVELOX (moxifloxacin hydrochloride) Tablets are available as oblong, dull red
- film-coated tablets containing 400 mg moxifloxacin. The tablet is coded with the
- word "BAYER" on one side and "M400" on the reverse side.

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831 Package

NDC Code

832 Bottles of 30:

0026-8581-69

833 ABC Pack of 5:

0026-8581-41

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- Store at 25°C (77°F); excursions permitted to 15-30°C (59-86°F) [see USP
- 836 Controlled Room Temperature]. Avoid high humidity.

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ANIMAL PHARMACOLOGY

- Quinolones have been shown to cause arthropathy in immature animals. In studies
- in juvenile dogs oral doses of moxifloxacin ≥ 30 mg/kg/day (approximately 1.5
- times the maximum recommended human dose based upon systemic exposure)
- for 28 days resulted in arthropathy. There was no evidence of arthropathy in
- mature monkeys and rats at oral doses up to 135 and 500 mg/kg, respectively.

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Unlike some other members of the quinolone class, crystalluria was not observed in 6 month repeat dose studies in rats and monkeys with moxifloxacin.

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- Ocular toxicity was not observed in 6 month repeat dose studies in rats and
- monkeys. In beagle dogs, electroretinographic (ERG) changes were observed in
- a 2 week study at doses of 60 and 90 mg/kg. Histopathological changes were
- observed in the retina from one of four dogs at 90 mg/kg, a dose associated with mortality in this study.

Some quinolones have been reported to have proconvulsant activity that is exacerbated with concomitant use of non-steroidal anti-inflammatory drugs (NSAIDS). Moxifloxacin at an oral dose of 300 mg/kg did not show an increase in acute toxicity or potential for CNS toxicity (e.g. seizures) in mice when used in combination with NSAIDs such as diclofenac, ibuprofen, or fenbufen.

In animal studies, at plasma concentrations about five times the human therapeutic level, a QT-prolonging effect of moxifloxacin was found. Electrophysiological *in vitro* studies suggested an inhibition of the rapid activating component of the delayed rectifier potassium current (I_{Kr}) as an underlying mechanism. In dogs, the combined infusion of sotalol, a Class III antiarrhythmic agent, with moxifloxacin induced a higher degree of QTc prolongation than that induced by the same dose (30mg/kg) of moxifloxacin alone.

CLINICAL STUDIES

Acute Bacterial Exacerbation of Chronic Bronchitis

AVELOX Tablets (400 mg once daily for five days) were evaluated for the treatment of acute bacterial exacerbation of chronic bronchitis in a large, randomized, double-blind, controlled clinical trial conducted in the US. This study compared AVELOX with clarithromycin (500 mg twice daily for 10 days) and enrolled 629 patients. The primary endpoint for this trial was clinical success at 7-17 days post-therapy. The clinical success for AVELOX was 89% (222/250) compared to 89% (224/251) for clarithromycin.

The following outcomes are the clinical success rates at the follow-up visit for the clinically evaluable patient groups by pathogen:

882	<u>PATHOGEN</u>	AVELOX	Clarithromycin
883			
884	Streptococcus pneumoniae	100% (16/16)	87% (20/23)
885	Haemophilus influenzae	89% (33/37)	88% (36/41)
886	Haemophilus parainfluenzae	100% (16/16)	100% (14/14)
887	Moraxella catarrhalis	85% (29/34)	100% (24/24)
888	Staphylococcus aureus	94% (15/16)	75% (6/8)
889	Klebsiella pneumoniae	90% (18/20)	91% (10/11)

The microbiological eradication rates (eradication plus presumed eradication) in AVELOX treated patients were *Streptococcus pneumoniae* 100%, *Haemophilus influenzae* 89%, *Haemophilus parainfluenzae* 100%, *Moraxella catarrhalis* 85%, *Staphylococcus aureus* 94%, and *Klebsiella pneumoniae* 85%.

Community Acquired Pneumonia

A large, randomized, double-blind, controlled clinical trial was conducted in the US to compare the efficacy of AVELOX Tablets (400 mg once daily) to that of high-dose clarithromycin (500 mg twice daily) in the treatment of patients with clinically

and radiologically documented community acquired pneumonia. This study enrolled 474 patients (382 of which were valid for the primary efficacy analysis conducted at the 14 - 35 day follow-up visit). Clinical success for clinically evaluable patients was 95% (184/194) for AVELOX and 95% (178/188) for high dose clarithromycin.

In addition to the trial described above, a noncomparative trial of AVELOX (400 mg once daily for ten days) was also conducted in the US in patients with community acquired pneumonia. The combined moxifloxacin clinical success rates by pathogen for the two studies were as follows:

910	<u>PATHOGEN</u>	<u>14 - 35 DAY FOLLOW-UP</u>
911		
912	Streptococcus pneumoniae	97% (30/31)
913	Haemophilus influenzae	92% (33/36)
914	Mycoplasma pneumoniae	96% (51/53)
915	Chlamydia pneumoniae	93% (106/114)
916	Moraxella catarrhalis	91% (10/11)

The microbiological eradication rates (eradication plus presumed eradication) in AVELOX treated patients were *Streptococcus pneumoniae* 97%, *Haemophilus influenzae* 92%, and *Moraxella catarrhalis* 91%.

Acute Bacterial Sinusitis

In a large, controlled double-blind study conducted in the US, AVELOX (400 mg once daily for ten days) was compared with cefuroxime axetil (250 mg twice daily for ten days) for the treatment of acute bacterial sinusitis. The trial included 457 patients valid for the primary efficacy determination. Clinical success (cure plus improvement) at the 7 to 21 day post-therapy test of cure visit was 90% for AVELOX and 89% for cefuroxime.

- An additional non-comparative study was conducted to gather bacteriological data and to evaluate microbiological eradication in adult patients treated with AVELOX 400 mg once daily for seven days. All patients (n = 336) underwent antral puncture in this study. Clinical success rates and eradication/presumed eradication rates at the 21 to 37 day follow-up visit were 97% (29 out of 30) for *Streptococcus pneumoniae*, 83% (15 out of 18) for *Moraxella catarrhalis*, and
- 936 80% (24 out of 30) for Haemophilus influenzae.

Uncomplicated Skin and Skin Structure Infections

A randomized, double-blind, controlled clinical trial conducted in the US compared the efficacy of AVELOX 400 mg once daily for seven days with Cephalexin HCl 500 mg three times daily for seven days. The percentage of patients treated for uncomplicated abscesses was 30%, furuncles 8%, cellulitis 16%, impetigo 20%, and other skin infections 26%. Adjunctive procedures (incision and drainage or debridement) were performed on 17% of the AVELOX treated patients and 14% of the comparator treated patients. Clinical success rates in evaluable patients were 89% (108/122) for AVELOX and 91% (110/121) for Cephalexin HCl.

REFERENCES 1. National Committee for Clinical Laboratory Standards, <u>Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria That Grow Aerobically-Fifth Edition</u>. Approved Standard NCCLS Document M7-A5, Vol. 20, No. 2, NCCLS, Wayne, PA, January 2000.

2. National Committee for Clinical Laboratory Standards, <u>Performance Standards for Antimicrobial Disk Susceptibility Tests</u>-Seventh Edition. Approved Standard NCCLS Document M2-A7, Vol. 20, No. 1, NCCLS, Wayne, PA, January, 2000.

Patient Information About:

AVELOX®

(moxifloxacin hydrochloride) 400 mg Tablets

This section contains important information about AVELOX (moxifloxacin hydrochloride), and should be read completely before you begin treatment. This section does not take the place of discussions with your doctor or health care professional about your medical condition or your treatment. This section does not list all benefits and risks of AVELOX. The medicine described here can be prescribed only by a licensed health care professional. If you have any questions about AVELOX talk with your health care professional. Only your health care professional can determine if AVELOX is right for you.

What is AVELOX?

 AVELOX is an antibiotic used to treat lung, sinus, or skin infections caused by certain germs called bacteria. AVELOX kills many of the types of bacteria that can infect the lungs and sinuses and has been shown in a large number of clinical trials to be safe and effective for the treatment of bacterial infections.

Sometimes viruses rather than bacteria may infect the lungs and sinuses (for example the common cold). AVELOX, like all other antibiotics, does not kill viruses.

You should contact your doctor if you think your condition is not improving while taking AVELOX. AVELOX Tablets are red and contain 400 mg of active drug.

How and when should I take AVELOX?

AVELOX should be taken once a day for 5, 7, or 10 days depending on your prescription. It should be swallowed and may be taken with or without food. Try to take the tablet at the same time each day.

You may begin to feel better quickly; however, in order to make sure that all bacteria are killed, you should complete the full course of medication. Do not take more than the prescribed dose of AVELOX even if you missed a dose by mistake. You should not take a double dose.

Who should not take AVELOX?

You should not take AVELOX if you have ever had a severe allergic reaction to any of the group of antibiotics known as "quinolones" such as ciprofloxacin or levofloxacin.

You should avoid AVELOX if you have a rare condition known as congenital prolongation of the QT interval. If you or any of your family members have this condition you should inform your health care professional. You should avoid AVELOX if you are being treated for heart rhythm disturbances with certain medicines such as quinidine, procainamide, amiodarone or sotalol. Inform your health care professional if you are taking a heart rhythm drug.

You should also avoid AVELOX if the amount of potassium in your blood is low. Low potassium can sometimes be caused by medicines called diuretics such as furosemide and hydrochlorothiazide. If you are taking a diuretic medicine you should speak with your health care professional.

If you are pregnant or planning to become pregnant while taking AVELOX, talk to your doctor before taking this medication. AVELOX is not recommended for use during pregnancy or nursing, as the effects on the unborn child or nursing infant are unknown.

AVELOX is not recommended for children.

What are the possible side effects of AVELOX?

- AVELOX is generally well tolerated. The most common side effects caused by
 AVELOX, which are usually mild, include nausea, vomiting, stomach pain,
 diarrhea, dizziness and headache. You should be careful about driving or
 operating machinery until you are sure AVELOX is not causing dizziness. If you
- notice any side effects not mentioned in this section or you have any concerns

about the side effects you are experiencing, please inform your health care 1028 professional. 1029 1030 In some people, AVELOX, as with some other antibiotics, may produce a small 1031 effect on the heart that is seen on an electrocardiogram test. Although this has 1032 not caused any serious problems in more than 4000 patients who have already 1033 1034 taken the medication in clinical studies, in theory it could result in extremely rare cases of abnormal heartbeat which may be dangerous. Contact your health care 1035 professional if you develop heart palpitations (fast beating), or have fainting spells. 1036 1037 Which medicines should not be used with AVELOX? 1038 1039 1040 You should avoid taking AVELOX with certain medicines used to treat an 1041 abnormal heartbeat. These include quinidine, procainamide, amiodarone, and sotalol. 1042 1043 1044 Some medicines also produce an effect on the electrocardiogram test, including cisapride, erythromycin, some antidepressants and some antipsychotic drugs. 1045 These may increase the risk of heart beat problems when taken with AVELOX. 1046 1047 For this reason it is important to let your health care provider know all of the medicines that you are using. 1048 1049 Many antacids and multivitamins may interfere with the absorption of AVELOX 1050 and may prevent it from working properly. You should take AVELOX either 4 1051 hours before or 8 hours after taking these products. 1052 1053 1054 Remember 1055 Take your dose of AVELOX once a day. 1056 1057 1058 Complete the course of medication even if you are feeling better. 1059 Keep this medication out of the reach of children. 1060 1061 This information does not take the place of discussions with your doctor or health 1062 1063 care professional about your medical condition or your treatment. 1064 **Bayer Corporation** 1065 Pharmaceutical Division 1066 400 Morgan Lane 1067 West Haven, CT 06516 1068 Made in Germany 1069 Rx Only 1070 **PZSKIN** 1071 04/01 ©2001 Bayer Corporation